PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

C12N 15/31, C07K 14/255, G01N 33/50

(11) International Publication Number:

WO 98/03656

(43) International Publication Date:

29 January 1998 (29.01.98)

(21) International Application Number:

PCT/US97/12639

A1

(22) International Filing Date:

18 July 1997 (18.07.97)

(30) Priority Data:

60/022,191

19 July 1996 (19.07.96)

US

(71) Applicant (for all designated States except US): REGENTS OF THE UNIVERSITY OF MINNESOTA [US/US]; Morrill Hall, 100 Church Street, S.E., Minneapolis, MN 55455 (US).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): RAJASHEKARA, Gireesh [IN/US]; Minn. Graduate Club, 2089 Carter Avenue, St. Paul, MN 55108 (US). NAGARAJA, Kakambi, V. [IN/US]; 2235 Milton Street North, Roseville, MN 55113 (US). KAPUR, Vivek [IN/US]; Apartment 108, 2572 Kenzie Terrace, St. Anthony, MN 55418 (US).
- (74) Agent: BRUESS, Steven, C.; Merchant, Gould, Smith, Edell, Welter & Schmidt, P.A., 3100 Norwest Center, 90 South Seventh Street, Minneapolis, MN 55402-4131 (US).

(81) Designated States: AL, AM, AT, AT (Utility model), AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, CZ (Utility model), DE, DE (Utility model), DK, DK (Utility model), EE, EE (Utility model), ES, FI, FI (Utility model), GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SK (Utility model), SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

- (54) Title: RECOMBINANT SEF14 FIMBRIAL PROTEIN FROM SALMONELLA
- (57) Abstract

A truncated SE fimbria antigen useful as an antigen for immunoassay diagnosis of Salmonella enteritidis (SE) infection or evidence of infection.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	. Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mati	· TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malewi	US	United States of America
CA	Canada	IT .	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
Cυ	Cuba	K2	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	u	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

20

RECOMBINANT SEF14 FIMBRIAL PROTEIN FROM SALMONELLA

Field of the Invention

The present invention relates to a method of cloning and expressing a truncated form of a fimbrial gene and the use of the truncated fimbrial gene product in an immunodiagnostic assay and for immunoprophylaxis.

Background of the Invention

10 Foodborne infections cause an estimated 6.5
million cases of human illness and 9000 deaths annually in
the United States alone. Bacterial infections by
Salmonella are the most commonly reported cause of
foodborne outbreaks. Salmonella enteritidis (SE) is the
15 dominant Salmonella serotype isolated from cases of food
poisoning. Many of these outbreaks are thought to be due
to infected poultry products, particularly eggs and egg
products.

The best way to prevent infection in human populations is to diagnose and treat the infected animal prior to human consumption. Because the greatest threat of food poisoning from Salmonella is from poultry products, there is a need for a method to detect birds that are infected with SE.

25 Some current diagnostic methods rely on conventional bacteriologic cultures. However, these procedures are relatively slow, often taking up to 3 to 4 days to provide even a presumptive diagnosis.

Additionally, the great susceptibility of SE to physical and chemical factors such as desiccation, radiation, low temperature, heating, or chemical preservatives, causes traditional bacteriologic culture methods to generally have a low sensitivity. Consequently, many birds or animals

10

15

20

25

30

that are infected with SE are often not detected when conventional bacterial cultures are used.

Other diagnostic methods rely on the detection of serum antibodies specific to SE. Although several serological methods such as micro-agglutination, serum plate agglutination, latex particle agglutination microantiglobulin, ELISA have previously been employed, these assays lack either the sensitivity or specificity necessary to detect SE infected birds, or the tests are too difficult to perform in a routine laboratory or field setting. Consequently, widespread application of these tests for the detection of SE infections has been impractical.

A useful antigenic determinant that is found on many species of Enterobacteriaceae are fimbriae, proteinaceous filamentous surface structures composed of protein subunits called fimbrin. Upon infection, birds make antibodies to this SE fimbrial antigen. Therefore, the SE fimbrial antigen is useful in a diagnostic assay for the presence of SE in poultry.

SE is known to have at least four distinct fimbria, designated Sef14, Sef17, Sef18 and Sef21. These proteins are encoded by SefA, AgfA, SefD and FinA genes, respectively.

Although the gene encoding Sef14 has been identified and its DNA nucleotide sequence determined (Trucotte and Woodward, Journal of General Microbiology, 139:1477-1485 (1993)), an effective diagnostic method using this surface antigen has not been developed, partially due to the difficulty of efficiently producing the fimbriae proteins in purified form and in large quantities. Additionally, expression of Sef14 fimbriae by cultured

10

15

20

25

Salmonella enteritidis is highly dependent on the growth medium composition. In a study by Thorns et al., International Journal of Food Microbiology, 21:47-53 (1994), only peptone water pH 7.2 supported the expression of Sef14 by all Salmonella enteritidis strains examined. Consequently, previous diagnostic assays using Sef14 have used antibodies against Sef14 and not the antigen itself.

Hence, there is a need for a sensitive, specific and routine antigen and method to reliably detect SE infection in birds, preferably a method that is easily adaptable to large-scale screening of poultry flocks.

Summary of the Invention

The present invention provides a sensitive, specific, routine antigen and assay to reliably detect SE-infected animals. Specifically, the present invention provides a truncated form of the Sef14 antigen that can be easily produced in purified form and in large quantities and used in the method of the invention. The novel Sef14 antigen, when coupled to a substrate such as latex beads, provides a diagnostic assay for SE, particularly useful in large-scale screening of poultry flocks.

Brief Description of the Figures

Figure 1 is a photograph showing a SDS-PAGE of the recombinant Sef14 (rSef14) fragment (arrow).

Figure 2 is a photograph showing a Western blot of the rSef14 fragment probed with anti-Sef14 antibody (lane 1) and anti-tag (T7) antibody (lane 2).

Figure 3 is a photograph showing results of a rSef14-latex agglutination assay for SE infection in

chickens exposed to S. enteritidis (A), S. pullorum (B), and serum-free antigen control (C).

Figure 4 is a photograph showing results of a rSef14-latex agglutination assay for SE infection in chickens exposed to S. enteritidis (A), S. gallinarum (B), S. pullorum (C), S. typhimurium (D), C. arizonae (E), E. coli (F), serum free antigen control (G), and serum control (H).

Figure 5 is a graph showing the percentage of chickens testing positive for anti-SE antibodies during 4 weeks post-innoculation. The five bars at each week represent innoculation with 10⁴, 10⁶, 10⁸, 10¹⁰, and control (no cells).

Figure 6 is a graph showing the antibody titres of chicken sera samples testing positive for anti-SE antibodies.

Figure 7 is a graph showing the antibody titres of chicken egg yolk samples testing positive for anti-SE antibodies.

20

25

30

5

10

15

Detailed Description of the Invention

The present invention is directed to a method for diagnosing Salmonella enteritidis infection or evidence of infection in an animal, particularly poultry, using a recombinant truncated fimbrial antigen.

"Infection" means active colonization of the animal by SE organisms. "Evidence of infection" means a prior history of colonization by SE in the animal, although active colonization is not present. Diagnosis of active infection is needed to protect against contamination of food supplies, whereas diagnosis of prior infection is

10

15

needed to alert against new infection or to trace the source of infection in a flock.

Fimbrial Proteins

Fimbriae are proteinaceous filamentous surface structures composed of protein subunits called fimbrin. These proteinaceous structures are thought to be virulence factors which mediate specific attachment to host cell mucosal surfaces. They are present in most enteric bacteria capable of invading host cells.

Salmonella enteritidis has four distinct fimbriae: Sef14, Sef17, Sef18 and Sef21 which are encoded by sefA, agfA, sefD and fimA genes, respectively. Sef14 is unique with only limited distribution in the genus. In contrast, all other fimbrial proteins are widely distributed in the genus. Thus, they have limited use as diagnostic reagents for SE detection.

Cloning and Expression of Sef14

In the present invention, a truncated form of the Sef14 antigen retaining the antigenic character of the entire protein has been produced. Unlike the complete protein, however, the truncated form can be easily produced in purified form and in large quantities, without special growth medium requirements.

PCR technology is used to produce the truncated Sef14 protein by amplification with suitable primers.

Primers are selected to amplify the gene encoding Sef14 in a region downstream of the encoded signal peptide, e.g., downstream of about nucleotide 145 of the DraI genomic fragment shown in Figure 1 of Turcotte and Woodward, Supra. Preferably, the PCR primers include

additional nucleotides at the 5' ends, encoding specific restriction enzyme recognition sequences, for ease of purification. For example, useful primers for amplifying that portion of the sefA gene encoding an immunogenic Sef14 fragment downstream of the signal peptide are shown below:

<u>GGGAATTC</u> GCTGGCTTTGTTGGTAACA	SEQ	ID	NO:1
GGGCTCGAGTTAGTTTTGATACTGAACGTA	SEQ	ID	NO:2

After a truncated gene sequence encoding Sef14 is produced, it can be cloned into a host using a plasmid or phage as a vector. Typically, the expression of Sef14 fimbriae by cultured Salmonella enteritidis is highly dependent on the growth medium composition (Thorns et al, International Journal of Food Microbiology, 21:47-53 (1994)), and it is typically difficult to produce large quantities. However, a truncated form of Sef14 having at least the signal peptide removed is expressed in host systems such as E. coli without these difficulties.

Truncated Sef14 Antigen

Because the truncated Sef14 protein retains the antigenic characteristics of the complete protein, it is useful in various immunological methods. For example, the inventive antigen is useful in antibody binding immunoassays such as assays to detect the presence of antibodies against SE in a sample. Suitable binding assays include ELISA, wherein the recombinant Sef14 antigen is bound to a surface and exposed to antibodies against SE. To detect the presence of bound anti-SE antibodies, a marker such as an enzyme-linked secondary antibody is then added.

5

10

An agglutination assay using truncated Sef14 antigen-coated latex beads is preferred. In the agglutination reaction, antigen-coated latex beads form detectable clusters when exposed to antibodies against SE. This preferred assay is described more fully in Example 4, below.

Diagnostic Assays

5

10

15

20

The assays described above can be used to detect the presence of antibodies to Salmonella enteritidis. Preferably, the assays are used to determine whether or not an animal, e.g. a poultry animal such as a chicken or turkey, is infected with SE. Animal fluid such as blood or serum can be used in a diagnostic assay. If an animal is infected with SE, the animal will typically produce anti-SE antibodies. The recombinant Sef14 antigen is used to detect the presence of anti-SE antibodies, SE infection or the SE organism itself. Diagnostic assays such as these are particularly useful in birds. More particularly, diagnostic assays are useful in detecting SE infections in chicken or turkey to prevent foodborne illness by poultry consumption.

Vaccine

Passive immunization with anti-Sef14 antibodies has been shown to reduce Salmonella enteritidis colonization (Peralta et al. 1994). Additionally, Sef14 can induce a T-cell immune response (Ogunniyi et al 1994). Because the truncated Sef14 antigen exhibits these immunological activities, can be produced in large quantities, and does not have the cumbersome growth requirements of the complete protein, the truncated Sef14

PCT/US97/12639

antigen is also useful as a vaccine to confer immunity against SE. Preferably, the truncated Sef14 antigen is used as a vaccine in poultry to prevent foodborne illnesses.

5

10

EXAMPLES

The invention may be better understood with reference to the following examples which are not intended to limit the invention.

Example 1

Isolation of S. enteritidis genomic DNA

S. enteritidis was grown overnight at 37°C in 15 Luria-Bertani (LB) broth. Genomic DNA was extracted as described (Sambrook, et al., 1989) using standard methods with minor modifications. In brief, bacterial cells were pelleted by centrifugation at 13,000 x g for 3 minutes, 20 washed/suspended in 1 ml of 1 M NaCl, centrifuged for 5 minutes at 13,000 x g, and the pellet resuspended in 1 ml TE buffer (50 mM Tris-HCl, 50 mM EDTA, pH 7.8). The sample was next incubated with 5 μl of lysozyme (50 mg/ml) (Sigma Chemical Co., St. Louis, MO) and 0.3 mg/ml RNase A (Sigma) at 37°C for 30 minutes. To this suspension, 1% sarkosyl 25 and 0.6 mg/ml of proteinase K (Sigma) were added, and the mixture incubated at 37° for 1 hour. Following incubation, chromosomal DNA was extracted twice with an equal volume of phenol-chloroform-isoamyl alcohol (25:24:1) and once with chloroform:isoamyl alcohol (24:1). Genomic DNA in the 30 aqueous phase was precipitated at -20°C with two volumes of absolute ethanol and 0.1 volume of 3 M sodium acetate, and

10

15

pelleted by centrifugation at 13,000 x g for 5 minutes. The pellet was then washed twice with 70% ethanol, air dried, and suspended in TE Buffer (10mM Tris-HCl and 1mM EDTA pH 8.0). Total DNA was quantitated spectrophotometrically at A_{250} nm.

Example 2

Cloning of sefA gene fragment

Oligonucleotide primer selection and synthesis:

Oligonucleotide primers corresponding to an internal fragment (64-498 bp) of the open-reading frame of the sefA gene were used for PCR amplification. Additional bases were added to the 5' end of each primer in order to confer a recognition sequence for either EcoRI (forward primer) or XhoI (reverse primer). The oligonucleotide primers were obtained from Integrated DNA technologies Inc., Ames, IA. The DNA sequences for the forward and reverse primers are shown below:

<u>GGGAATTC</u> GCTGGCTTTGTTGGTAACA	SEQ	ID	NO:1
GGGCTCGAGTTAGTTTTGATACTGAACGTA	SEQ	ID	NO:2

20 Additional nucleotides added to the 5' end of the primers are underlined.

PCR amplification of sefA gene fragment.

Amplification reactions were performed in 30 μ l volumes with 30 pmol of each primer and 5 mM MgCl₂. The reagents and enzymes used for PCR were obtained either from Boehringer Mannheim (Indianapolis, IN) or Perkin Elmer (Foster City, CA). One hundred ng of genomic DNA was used as a template for PCR amplification with the following parameters: an initial denaturation at 94°C for 5 minutes, followed by 35 cycles of denaturation (94°C for 1.5

25

minutes), annealing 52°C for 1 minute) and extension (72°C for 2 minutes), and a final extension of 15 minutes at 72°C. All amplification reactions were performed in a Perkin-Elmer Cetus DNA thermal cycler (Model 480). The PCR products were analyzed on a 1% agarose gel, stained with ethidium bromide (0.5µg/ml), and photographed under UV light.

PCR products were gel extracted (Qiagen Inc., Chatsworth, CA), quantitated spectophometrically, at 260 nm, and cloned directly into pGEM-T vector (Promega, Madison, WI). Following ligation, 2µl of the reaction products were transformed into E. coli DH5α cells (Gibco BRL, Gaithersburg, MD) by the heat shock method. Recombinant colonies were selected on ampicillin/IPTG-Xgal containing plates and screened for the presence of the appropriate insert by restriction analysis.

Nucleotide sequence analysis

A bacterial colony containing the recombinant plasmid with the rSefA fragment was grown in LB-ampicillin media, and the plasmid extracted using Qiagen plasmid extraction kit (Qiagen). The nucleotide sequence of the insert was determined using oligonucleotide primers specific to the vector sequence by automated DNA sequencing at the University of Minnesota Advanced Genetic Analysis 25 The insert was sequenced in its entirety in both orientations, and the amino acid sequence deduced using the standard genetic code (DNA*, Madison, WI). Sequencing results are shown below for nucleotide and deduced amino acid sequences of the insert (Seq.ID.NO:5), together with a 30 tag sequence added during the subcloning of the fragment into the pET/abc expression vector (Seq.ID.NO:3).

5

10

15

added tag sequence at the 5' end, provides a Histidine-rich portion to facilitate purification of the sequence on nickel columns, as well as an antigenic region that specifically binds the T7 anti-tag antibody provided with the pET/abc vector kit.

Nucleic Acid Sequence encoding rSefA fragment SEQ ID NO:3

ATG GGC AGC AGC CAT CAT CAT CAT CAC AGC AGC GGC CTG GTG 45 CCG CGC GGC AGC CAT ATG GCT AGC ATG ACT GGT GGA CAG CAA ATG 90 GGT CGC GGA TGG GAA TTC GCT GGC TTT GTT GGT AAC AAA GCA GTG GTT CAG GCA GCG GTT ACT ATT GCA GCT CAG AAT ACA ACA TCA GCC AAC TGG AGT CAG GAT CCT GGC TTT ACA GGG CCT GCT GCT GCT 225 GGT CAG AAA GTT GGT ACT CTC AGC ATT ACT GCT ACT GGT CCA CAT 270 AAC TCA GTA TCT ATT GCA GGT AAA GGG GCT TCG GTA TCT GGT GGT 315 GTA GCC ACT GTC CCG TTC GTT GAT GGA CAA GGA CAG CCT GTT TTC 360 CGT GGG CGT ATT CAG GGA GCC AAT ATT AAT GAC CAA GCA AAT ACT 405 GGA ATT GAC GGG CTT GCA GGT TGG CGA GTT GCC AGC TCT CAA GAA 450 ACG CTA AAT GTC CCT GTC ACA ACC TTT GGT AAA TCG ACC CTG CCA 495 GCA GGT ACT TTC ACT GCG ACC TTC TAC GTT CAG CAG TAT CAA AAC 540 TAA CTC GAG CCC 552

*Additional amino acid residues added to the amino terminus to facilitate protein purification and cloning are underlined.

Deduced amino acid sequence of rSefA protein fragment*

Seq. ID NO: 4

MGSSHHHHHHSSGLVPRGSHMASMTGGOOMGRGSEFAGFVGNKAVVQAAVT

IAAQNTTSANWSQDPGFTGPAVAAGQKVGTLSITATGPHNSVSIAGKGASVSGG

VATVPFVDGQGQPVFRGRIQGANINDQANTGIDGLAGWRVASSQETLNVPVTT

FGKSTLPAGTFTATFYVQQYQN

*Additional amino acid residues added to the amino terminus to facilitate protein purification and cloning are underlined.

It is understood that the amino acids added to the N-terminus of the Spf14 antigen are optional, and used for ease of cloning and purification. The amino acid sequence in the absence of these added residues (Sequence ID No:6)

15

20

with or without other added residues for cloning or purification procedures, for example, are similarly useful as antigens in the diagnostic assays of the invention.

Subcloning sefA gene fragment into an expression vector

The pGEM-T plasmid carrying sefA fragment was double digested with EcoRI and XhoI, and the digested products gel purified (Qiagen) and cloned into EcoRI and XhoI digested pET/abc expression vectors (Novagen Inc., Madison, WI). Ligation products (2 µl) from each of the reactions were transformed into E. coli BL21(DE3)pLyS cells by heat shock method. Recombinant clones were cultured on kanamycin and chloramphenicol containing plates, and analyzed by restriction enzyme analysis.

15

20

25

30

10

5

rSefA fragment expression

The recombinant clones were selected based on restriction enzyme analysis with EcoRI and XhoI digestion, selecting those clones yielding appropriately sized fragments as compared with a vector control. Selected clones were analyzed for rSefA fragment expression. Briefly, a single colony from each (pETabc/SefA fragment) freshly streaked plate was picked and inoculated to 50 ml LB broth containing appropriate antibiotics and incubated with shaking at 200 rpm at 37°C until the OD₆₀₀ reached 0.6. Cultures were induced with IPTG (0.4 mM) and incubated for an additional 3 hours. Following incubation, the cells were pelleted and resuspended in 5 ml of TE buffer (50mM Tris-HCl pH 8.0, 2mM EDTA) and incubated with 25 µl of lysozyme (50 mg/ml) and 100 μ l of 1% Triton X-100 for 20 minutes at 30°C. The samples were sonicated until they were no longer viscous, and centrifuged at 39,000 x g for

20 minutes. The supernatant was passed through a 0.45 μm membrane filter, and stored at -20°C until further use.

SDS-PAGE analysis

5

10

The cell lysates were next analyzed by SDS-PAGE for the presence of the rSefA fragment by mixing with an equal volume of 2x SDS solubilization buffer separating on 12% polyacrylamide gels, and staining with Coomassie blue. The results are shown in lane 1 of Figure 1 which contains the total protein produced by the vector and contained in the cell lysates.

Western blot analysis

The lysates were separated on 12% polyacrylamide 15 gels and transferred onto a nitrocellulose membrane using Transblot apparatus (Bio-Rad laboratories, Hercules, CA). Following transfer, the membrane was blocked with 3% BSA in phosphate buffered saline (PBS) and stained with either T7 anti-tag antibody (Novagen) or rabbit anti-Sef14 specific 20 antibody (kindly provided by Dr. W. W. Kay, University of Victoria, BC, Canada). The membrane was washed and stained with anti-rabbit IgG/HRP conjugate and treated with developing reagent (Amersham lif sciences, Inc., USA) for 1 minute, exposed to X-ray film, and the radiograph developed. The results are shown in Figure 2, where lane 1 25 is probed with anti-Sef14 antibody, and lane 2 with T7 anti-tag antibody.

Purification of rSef14 fragment protein by column chromatography and electroelution

The recombinant Sef14 protein fragment produced in the cell lysates described above was purified by binding

of the Histidine-rich tag to nickel columns as described by the manufacturer (Novagen). Briefly, the cells were induced and extract was prepared as described above except that the induced cells were suspended in Tris buffer without EDTA. The cell lysate was passed through nickel 5 columns and washed sequentially with binding buffer (5 mM imidazole, 0.5 M NaCl, 20 mM Tris-CHl, pH 7.9) and wash buffer (60 mM imidazole, 0.5 M NaCl, 20 mM Tris-HCl, pH The bound protein was eluted using elution buffer (1 M imidazole, 0.5 M NaCl, 20 mM Tris-HCl, pH 7.9), 10 quantitated using a Bio-Rad protein assay kit (Bio-Rad laboratories, Hercules, CA), and analyzed by SDS-PAGE. (See Figure 1, lane 2, where the arrow indicates the rSef14 fragment at about 19 KDa.) Since the column purified recombinant material contained traces of non-specific 15 proteins, the appropriate rSef14 fragment was further purified by cutting the rSef14 fragment from the gel and electroelution (Bio-Rad) following the manufacturer's suggested protocol. The electroluted fragment is shown in lane 4 of Figure 1 (at arrow). 20

Example 3

Covalent coupling of rSef14 to blue-dyed latex beads

The electroeluted rSef14 protein fragment was coupled to either 0.5 µm or 1.0 µm blue-dyed latex beads (Polysciences Inc., Warrington, PA) by gluteraldehyde method. Briefly, 1 ml of 2.5% suspension of the beads were washed with PBS (pH 7.4), pelleted by centrifugation and resuspended in 1 ml of 8% gluteraldehyde (EM grade) in PBS, and incubated overnight with gentle end-to-end mixing at room temperature. Following gluteraldehyde treatment, the beads were pelleted, washed with PBS three times and

25

incubated with 500 μg of purified rSef14 fragment for 5 hours at room temperature with gentle end-to-end mixing. The beads were pelleted, and incubated with 1 ml of 0.5 M ethanolamine in PBS for 30 minutes at room temperature with gentle end-to-end mixing. The mixture was then treated with 1 ml of 10 mg/ml BSA in PBS for 30 minutes at room temperature, centrifuged and the pellet resuspended in 1 ml PBS (pH 7.4), containing 10 mg/ml BSA, 0.1% NaN₃ and 5% glycerol, and stored at 4°C to form rSef14 - fragment coated latex beads for use in agglutination assays.

5

10

BNSDOCID: <WO__9803656A1_1 >

Example 4

rSef14-latex bead agglutination test

Bacteria was administered to chickens by either injection, intratracheal or oral administration of 107 15 colony forming units (CFU) of either S. enteritidis, S. pullorum, S. arizonae, S. typhimurium, S. gallinarum, or E. coli. After about two to three weeks exposure, serum was collected and used to evaluate the sensitivity and 20 specificity of the rSef14-latex beads in an agglutination assay for anti-SE antibody binding. A total volume of 7.5 μl of rSef14 fragment coated latex beads, produced as described for Example 3, were mixed with an equal volume of chicken serum collected from birds exposed to various pathogens, as described above. The presence of 25 agglutination, visually seen as a loss of intense blue color in the sample (i.e., lightening of color as the coated beads agglutinate or form a lattice). Absence of the applutination reaction was visualized by the remaining intense blue color of the dyed beads in a homogeneous 30 suspension. Positive or negative agglutination reaction

10

15

20

25

30

was recorded after two minutes. The results are shown in Figures 3 and 4.

In figure 3, intense blue color (negative result) is seen in test samples B and C (S. pullorum and the serum-free antigen control). In contrast, a positive agglutination result is seen in test Sample A, (S. enteritidis), as a pale blue, diffuse agglutination pattern.

In figure 4, a positive agglutination reaction is seen in sample A (S. enteritidis) and in sample H (serum control). No agglutination reaction is seen in the samples B-G containing serum animals exposed to the following pathogens: S. gallinarium (B), S. pullorum (C), S. typhimurium (D), S. arizonae (E), E. coli (F), and serum free antigen control (H).

Example 5

Detection of anti-S.E. anithodies in infected chickens

To confirm the specificity of the assay of the invention, forty SPF chickens (age 4weeks) were innoculated with various species of Salmonella. A suspension of 10° CFU in PBS was administered by injection. A booster dose of 10° CFU was administered orally two weeks later. Serum samples were taken at weekly intervals and assayed for the presence of anti-SE antibodies.

5

10

15

20

BNSDOCID: <WO__9803656A1_T_

two assays are standard screening methods for the detection of Salmonella, using *S.pullorium* as a whole-cell antigen, and are not specific for SE, as shown in the table below.

17

To demonstrate the specificity of the assays of the invention, serum samples were assayed using the latex agglutination test (LAT) described above for Example 4, which utilized the truncated Spf14 antigen coupled to latex beads. Serum samples were also assayed for anti-SE antibodies by ELISA. In the ELISA, the truncated Spf14 antigen prepared as described for Example 3, was coated onto polystyrine plates. Antigen-coated plates were exposed to serum samples to permit binding of anti-SE antibodies to the antigen. The bound antigen-antibody complexes were washed, and then incubated with anti-chicken antibody coupled to biotin. The complex was then exposed to strep-avidin for signal detection.

Results are shown in the table below. The LAT and Elisa assays demonstrated a useful specificity for the detection of SE. Of the organisms tested, only *S.dublin*, a bovine pathogen, demonstrated cross-reactivity in the assays.

	Species	SPT	MT	LAT	ELISA
S.	enteritidis	+	+	+	+
S.	gallinarum	+	+	•	-
s.	pullorum	+	+	-	-
s.	dublin	+	+	+	+
s.	berta	+	+	-	-
s.	typhimurium	-	+	-	-
E.	coli	-	-	-	-
Co	ntrol (no cells)	-	-	-	-

Example 6

Specificity of anti-SE assay

The ELISA assay for detecting anti-SE antibodies described above for Example 6 was tested for specificity using a panel of antisera against known pathogenic organisms. Each sera was assayed in the anti-SE ELISA. No crossreativity was observed with any of the tested antisera.

10

5

Antisera	ELISA		Antisera	ELISA
Pox	-	77	MG	-
Reo	-		NDV	-
Rev	-		CAV	_
SB-1	-		HVT	
IBDV	-	12	IBV	-
ILT	-	** **	S.typhimurum	-
LLA	-		S.gallinarum	_
LLB	-		S.pullorum	-
MS	-	*		

15

20

Example 7

Sensitivity of ELISA for detection of SE

Fifty white leghorn layer chickens (5 weeks old) were orally innoculated in a single exposure with varied amounts of SE, from 10^4 to 10^{10} CFU in PBS. Serum samples were collected at weekly intervals for up to seven weeks. Eggs were collected for egg yolk antibody detection.

Samples were analized for detection of anti-SE

19

antibodies using the ELISA described above for Example 6. As shown in Figure 5, control chickens showed no positive reaction in the ELISA assay. Approximately 40-80% of chickens exposed to 10⁴, 10⁶, 10⁸, and 10¹⁰ CFU of SE tested positive for anti-SE antibodies during the first four weeks post-innoculation. From 4-7 weeks post-innoculation, the data stabilized at about 45% positive detection of anti-SE antibodies.

Antibody titers in the sera and egg yolks of chickens exposed to 10^4 , 10^6 , and 10^8 CFU of SE and testing positive in the ELISA for anti-SE antibodies are shown in Figures 6 and 7.

These data demonstrate specific detection of anti-SE antibodies using recombinant Sef14-antigen coated latex beads in an agglutination assay and using the antigen as a capture agent in an ELISA. These assays provide a sensitive and specific diagnostic tool for the detection of anti-SE antibodies in animals and for the diagnosis of SE infection.

5

10

SEQUENCE LISTING

- (1) GENERAL INFORMATION
- (i) APPLICANT: REGENTS OF THE UNIVERSITY OF MINNESOTA
- (ii) TITLE OF THE INVENTION:

 RECOMBINANT FIMBRIAL PROTEIN
- (iii) NUMBER OF SEQUENCES: 6
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Merchant, Gould, Smith, Edell, Welter & Schmidt
 - (B) STREET: 3100 Norwest Center, 90 South Seventh St
 - (C) CITY: Minneapolis
 - (D) STATE: MN
 - (E) COUNTRY: USA
 - (F) ZIP: 55402
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Diskette
 - (B) COMPUTER: IBM Compatible
 - (C) OPERATING SYSTEM: DOS
 - (D) SOFTWARE: FastSEQ Version 2.0
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE: 18-JUL-1997
 - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: 60/022,191
 - (B) FILING DATE: 19-JUL-1996
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Kettelberger, Denise M
 - (B) REGISTRATION NUMBER: 33,924
 - (C) REFERENCE/DOCKET NUMBER: 600.335W001
- (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 612/371-5268
 - (B) TELEFAX: 612/332-9081
 - (C) TELEX:

(2) INFORMATION FOR SEQ ID NO:1:

(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 27 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear
(ii) MOLECULE TYPE: cDNA
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
GGGAATTCGC TGGCTTTGTT GGTAACA 2
(2) INFORMATION FOR SEQ ID NO:2:
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 30 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear
(ii) MOLECULE TYPE: cDNA
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
GGGCTCGAGT TAGTTTTGAT ACTGAACGTA
(2) INFORMATION FOR SEQ ID NO:3:
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 552 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
(ii) MOLECULE TYPE: cDNA (ix) FEATURE:
(A) NAME/KEY: Coding Sequence (B) LOCATION: 1540 (D) OTHER INFORMATION:
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:
ATG GGC AGC AGC CAT CAT CAT CAT CAC AGC AGC GGC CTG GTG CCG Met Gly Ser Ser His His His His His Ser Ser Gly Leu Val Pro 1 5 10 15
CGC GGC AGC CAT ATG GCT AGC ATG ACT GGT GGA CAG CAA ATG GGT CGC Arg Gly Ser His Met Ala Ser Met Thr Gly Gly Gln Gln Met Gly Arg 20 25 30

				GGC Gly									144
				GCT Ala									192
				GGG Gly 70									240
	-			GCT Ala									288
				GTA Val									336
				CCT Pro			_	-			_		384
	-			AAT Asn									432
-				GAA Glu 150									480
				GCA Ala								CAG Gln	528
	TAT Tyr		TAAC	CTCGA	AGC (ec							552

(2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 180 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
 (v) FRAGMENT TYPE: internal
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met 1	Gly	Ser	Ser	His 5	His	His	His	His	His 10	Ser	Ser	Gly	Leu	Val 15	Pro
Arg	Gly	Ser	His 20	Met	Ala	Ser	Met	Thr 25	Gly	Gly	Gln	Gln	Met 30	Gly	Arg
Gly	Trp	Glu 35	Phe	Ala	Gly	Phe	Val 40	Gly	Asn	Lys	Ala	Val 45	Val	Gln	Ala
Ala	Val 50	Thr	Ile	Ala	Ala	Gln 55	Asn	Thr	Thr	Ser	Ala 60	Asn	Trp	Ser	Gln
Asp 65	Pro	Gly	Phe		Gly .70	Pro	Ala	Val	Ala	Ala 75	Gly	Gln	Ļys	Val	Gly 80
Thr	Leu	Ser	Ile	Thr 85	Ala	Thr	Gly	Pro	His 90	Asn	Ser	Val	Ser	Ile 95	Ala
Gly	Lys	Gly	Ala 100	Ser	Val	Ser	Gly	Gly 105	Val	Ala	Thr	Val	Pro 110	Phe	Val
Asp	Gly	Gln 115	Gly	Gln	Pro	Val	Phe 120	Arg	Gly	Arg	Ile	Gln 125	Gly	Ala	Asn
Ile	Asn 130	Asp	Gln	Ala	Asn	Thr 135	Gly	Ile	Asp	_	Leu 140	Ala	Gly	Trp	Arg
Val 145	Ala	Ser	Ser	Gln	Glu 150	Thr	Leu	Asn	Val	Pro 155	Val	Thr	Thr	Phe	Gly 160
Lys	Ser	Thr	Leu	Pro 165	Ala	Gly	Thr	Phe	Thr 170	Ala	Thr	Phe	Tyr	Val 175	Gln
Gln	Tyr	Gln	Asn 180												

- (2) INFORMATION FOR SEQ ID NO:5:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 435 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Genomic DNA
- (ix) FEATURE:
 - (A) NAME/KEY: Coding Sequence
 - (B) LOCATION: 1...432
 - (D) OTHER INFORMATION:
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

							ACT Thr 15	48
 	 						GGC Gly	96
 	 						AGC Ser	144

24

	GCT Ala 50									192
	GTA Val									240
	CCT Pro									288
	AAT Asn									336
	GAA Glu									384
	GCA Ala 130							AAC Asn	T	433
ממ							•			435

(2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 144 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (v) FRAGMENT TYPE: internal
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

10

WE CLAIM:

1. A method for detecting anti- Salmonella enteritidis antibodies in animals, the method comprising:

reacting a sample obtained from an animal with a truncated Sef14 antigen under conditions to permit anti-Sef14 antibodies to bind the antigen, the truncated antigen having at least the native Sef14 signal peptide removed; and

correlating antibody-antigen binding with the presence of anti-SE antibodies in the sample.

- 2. A method for diagnosing Salmonella enteritidis infection in animals, the method comprising:
- reacting a sample obtained from an animal with a truncated Sef14 antigen under conditions to permit anti-Sef14 antibodies to bind the antigen, the truncated antigen having at least the native Sef14 signal peptide removed; and
- 20 correlating antibody-antigen binding with Salmonella enteritidis infection.
 - 3. The method of claim 1, wherein the truncated Sef14 antigen has the amino acid sequence of Sequence ID. No. 4.
 - 4. The method of claim 1, wherein the truncated Sef14 antigen has the amino acid sequence of Sequence ID. No.6.
- 5. The method of claim 1, wherein said antigen is fixed to an inert surface prior to said reacting.

6. A Sef14 antigen consisting essentially of the amino acid sequence of Sequence I.D. No. 6.

- 7. An assay kit for the detection of anti-Salmonella enteritidis antibodies comprising an Sef14 antigen consisting essentially of the amino acid sequence of Sequence ID No. 6.
- 8. The assay kit of claim 7, wherein the antigen consists
 10 essentially of the amino acid sequence of Sequence ID. No.
 4.
 - 9. An antigen for stimulating the production of antiSalmonella enteritidis antibodies comprising the amino acid
 sequence of Sequence ID No. 4 or 6.
 - 10. The method of detecting anti-Salmonella enteritidis antibodies described in any of the foregoing claims, wherein the animal samples are obtained from fowl, and particularly from chickens or turkeys.

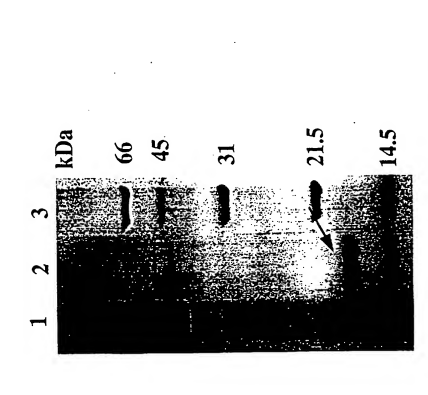
5

15

rSef14

Figure 1

SDS-PAGE of rSef14 fragment protein



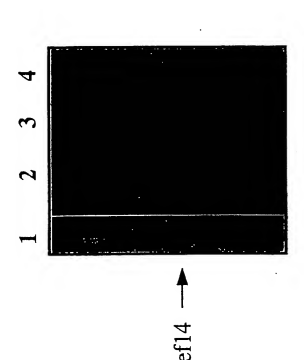
3. LMW marker

4. rSef14 after electroelution

BL21DE3 (pET/sefA)before purification
 Purified rSef14 fragment protein

HIGMIC C

Western blot -- rSef14 fragment probed with Sef14 monospecific polyclonal antibody



Anti-Sef14
 T7 tag antibody
 BL21DE3 control
 LMW marker

Figure 2

Later agglutination test

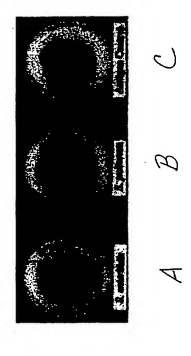


Figure 4

1. Latex agglutination test.

E. SDOCID: <WO__9803656A1_I_>

5/7.

Figure 5

Sensitivity of rSEF14-LAT Chicken serum

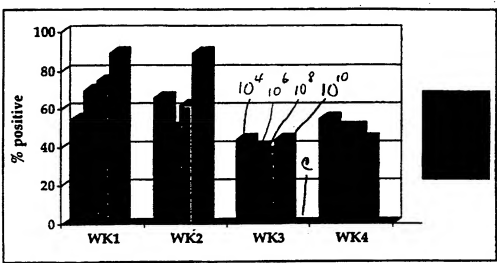
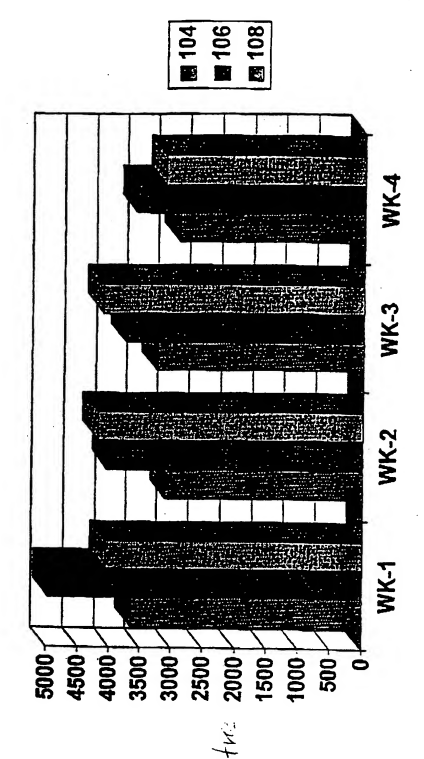
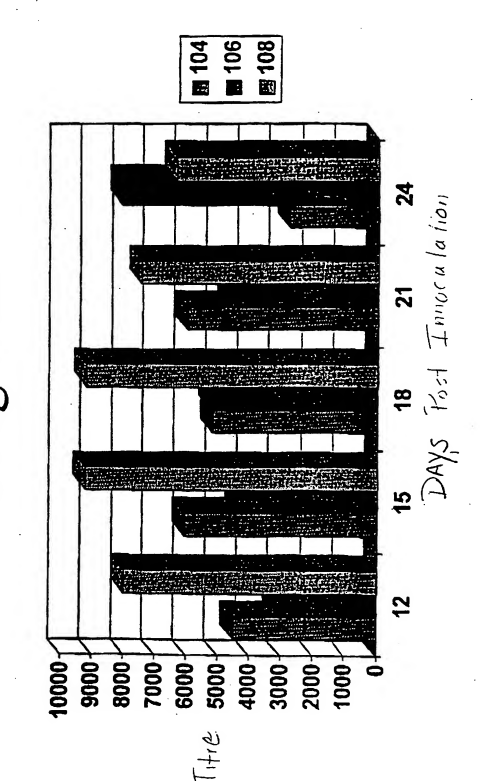


Figure 6

Sensitivity of SEF-14 ELISA Using Chicken Sera



Sensitivity of SEF-14 ELISA Using Yoll tigure 7



INTERNATIONAL SEARCH REPORT

into onal Application No PCT/US 97/12639

CLASSIFICATION OF SUBJECT MATTER IPC 6 CO7K14/255 G01N33/50 C12N15/31 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) C12N CO7K GO1N IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Category ' Citation of document, with indication, where appropriate, of the relevant passages THORNS CJ ET AL: "Development and 1,2,5,6, X application of enzyme-linked immunosorbent assay for specific detection of Salmonella enteritidis infections in chickens based on antibodies to SEF14 fimbrial antigen." J CLIN MICROBIOL, APR 1996, 34 (4) P792-7, UNITED STATES, XP002047275 see page 792 - page 793; table 1 1,2,5,6, WO 92 06197 A (MINI AGRICULTURE & Х FISHERIES) 16 April 1992 see claims 2.3 -/--Patent family members are listed in annex. X Further documents are listed in the continuation of box C. X * Special categories of cited documents : T later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) You document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled in the art. other means *P* document published prior to the international filing data but later than the priority data claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 0 2. 12. 97 18 November 1997 **Authorized officer** Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Espen, J

Form PCT/ISA/210 (second sheet) (July 1992)

Intern: al Application No PCT/US 97/12639

Continu	Ition) DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/US 97/	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	F	elevant to claim No.
Y	THORNS CJ ET AL: "The use of latex particle agglutination to specifically detect Salmonella enteritidis." INT J FOOD MICROBIOL, JAN 1994, 21 (1-2) P47-53, NETHERLANDS, XP002047276 see the whole document		1,2,5,6, 10
,	WO 92 06198 A (MINI AGRICULTURE & FISHERIES) 16 April 1992 see claims 1-28		1,2,5,6, 10
'	WO 93 20231 A (MINI AGRICULTURE & FISHERIES; WOODWARD MARTIN JOHN (GB); THORNS CH) 14 October 1993 see claims 1-28		1,2,5,6, 10
A	CLOUTHIER SC ET AL: "Characterization of three fimbrial genes, sefABC, of Salmonella enteritidis." J BACTERIOL, MAY 1993, 175 (9) P2523-33, UNITED STATES, XP002047277		
			•
			•

1

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

information on patent family members

Inter and Application No
PCT/US 97/12639

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 9206197 A	16-04-92	AU 660152 B	15-06-95
NO 3200137 A	10 04 52	AU 8548991 A	28-04-92
		CA 2091982 A	02-04-92
		EP 0551325 A	21-07-93
		JP 6501934 T	03-03-94
		US 5510241 A	23-04-96
	-	AT- 155169 T	15-07-97
		AU 660945 B	13-07-95
		AU 8656691 A	28-04-92
-	•	CA 2091984 A	02-04-92
		DE 69126786 D	14-08-97
	•	EP 0551324 A	21-07-93
		WO 9206198 A	16-04-92
		JP 6502531 T	24-03-94
WO 9206198 A	16-04-92	AT 155169 T	15-07-97
NO 3200230 //	20 0 1 72	AU 660945 B	13-07-95
_	•	AU 8656691 A	28-04-92
		CA 2091984 A	02-04-92
		DE 69126786 D	14-08-97
		EP 0551324 A	21-07-93
		JP 6502531 T	24-03-94
		AU 660152 B	15-06-95
		AU 8548991 A	28-04-92
		CA 2091982 A	02-04-92
		EP 0551325 A	21-07-93
	•	WO 9206197 A	16-04-92
		JP 6501934 T	03-03-94
		US 5510241 A	23-04-96
WO 9320231 A	14-10-93	AU 3895293 A	08-11-93



WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6: C12N 15/31, C07K 14/255, G01N 33/50

(11) International Publication Number:

WO 98/03656

A1 |

(43) International Publication Date:

29 January 1998 (29.01.98)

(21) International Application Number:

PCT/US97/12639

(22) International Filing Date:

18 July 1997 (18.07.97)

(30) Priority Data:

60/022,191

19 July 1996 (19.07.96)

US

(71) Applicant (for all designated States except US): REGENTS OF THE UNIVERSITY OF MINNESOTA [US/US]; Morrill Hall, 100 Church Street, S.E., Minneapolis, MN 55455 (US).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): RAJASHEKARA, Gireesh [IN/US]; Minn. Graduate Club, 2089 Carter Avenue, St. Paul, MN 55108 (US). NAGARAJA, Kakambi, V. [IN/US]; 2235 Milton Street North, Roseville, MN 55113 (US). KAPUR, Vivek [IN/US]; Apartment 108, 2572 Kenzie Terrace, St. Anthony, MN 55418 (US).
- (74) Agent: BRUESS, Steven, C.; Merchant, Gould, Smith, Edell, Welter & Schmidt, P.A., 3100 Norwest Center, 90 South Seventh Street, Minneapolis, MN 55402-4131 (US).

(81) Designated States: AL, AM, AT, AT (Utility model), AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, CZ (Utility model), DE, DE (Utility model), DK, DK (Utility model), EE, EE (Utility model), ES, FI, FI (Utility model), GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SK (Utility model), SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: RECOMBINANT SEF14 FIMBRIAL PROTEIN FROM SALMONELLA

(57) Abstract

A truncated SE fimbria antigen useful as an antigen for immunoassay diagnosis of Salmonella enteritidis (SE) infection or evidence of infection.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
. AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
- BA	Bosnia and Herzegovina	GE	Georgia	. MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	ĦU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	TI	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China .	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan .		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore -		

RECOMBINANT SEF14 FIMBRIAL PROTEIN FROM SALMONELLA

Field of the Invention

The present invention relates to a method of

cloning and expressing a truncated form of a fimbrial gene
and the use of the truncated fimbrial gene product in an
immunodiagnostic assay and for immunoprophylaxis.

Background of the Invention

10 Foodborne infections cause an estimated 6.5
million cases of human illness and 9000 deaths annually in
the United States alone. Bacterial infections by
Salmonella are the most commonly reported cause of
foodborne outbreaks. Salmonella enteritidis (SE) is the
dominant Salmonella serotype isolated from cases of food
poisoning. Many of these outbreaks are thought to be due
to infected poultry products, particularly eggs and egg
products.

The best way to prevent infection in human populations is to diagnose and treat the infected animal prior to human consumption. Because the greatest threat of food poisoning from Salmonella is from poultry products, there is a need for a method to detect birds that are infected with SE.

Some current diagnostic methods rely on conventional bacteriologic cultures. However, these procedures are relatively slow, often taking up to 3 to 4 days to provide even a presumptive diagnosis.

Additionally, the great susceptibility of SE to physical and chemical factors such as desiccation, radiation, low temperature, heating, or chemical preservatives, causes traditional bacteriologic culture methods to generally have a low sensitivity. Consequently, many birds or animals

10

15

20

25

30

that are infected with SE are often not detected when conventional bacterial cultures are used.

Other diagnostic methods rely on the detection of serum antibodies specific to SE. Although several serological methods such as micro-agglutination, serum plate agglutination, latex particle agglutination microantiglobulin, ELISA have previously been employed, these assays lack either the sensitivity or specificity necessary to detect SE infected birds, or the tests are too difficult to perform in a routine laboratory or field setting. Consequently, widespread application of these tests for the detection of SE infections has been impractical.

A useful antigenic determinant that is found on many species of *Enterobacteriaceae* are fimbriae, proteinaceous filamentous surface structures composed of protein subunits called fimbrin. Upon infection, birds make antibodies to this SE fimbrial antigen. Therefore, the SE fimbrial antigen is useful in a diagnostic assay for the presence of SE in poultry.

SE is known to have at least four distinct fimbria, designated Sef14, Sef17, Sef18 and Sef21. These proteins are encoded by SefA, AgfA, SefD and FinA genes, respectively.

Although the gene encoding Sef14 has been identified and its DNA nucleotide sequence determined (Trucotte and Woodward, Journal of General Microbiology, 139:1477-1485 (1993)), an effective diagnostic method using this surface antigen has not been developed, partially due to the difficulty of efficiently producing the fimbriae proteins in purified form and in large quantities.

Additionally, expression of Sef14 fimbriae by cultured

10

15

20

25

30

Salmonella enteritidis is highly dependent on the growth medium composition. In a study by Thorns et al., International Journal of Food Microbiology, 21:47-53 (1994), only peptone water pH 7.2 supported the expression of Sef14 by all Salmonella enteritidis strains examined. Consequently, previous diagnostic assays using Sef14 have used antibodies against Sef14 and not the antigen itself.

Hence, there is a need for a sensitive, specific and routine antigen and method to reliably detect SE infection in birds, preferably a method that is easily adaptable to large-scale screening of poultry flocks.

Summary of the Invention

The present invention provides a sensitive, specific, routine antigen and assay to reliably detect SE-infected animals. Specifically, the present invention provides a truncated form of the Sef14 antigen that can be easily produced in purified form and in large quantities and used in the method of the invention. The novel Sef14 antigen, when coupled to a substrate such as latex beads, provides a diagnostic assay for SE, particularly useful in large-scale screening of poultry flocks.

Brief Description of the Figures

Figure 1 is a photograph showing a SDS-PAGE of the recombinant Sef14 (rSef14) fragment (arrow).

Figure 2 is a photograph showing a Western blot of the rSef14 fragment probed with anti-Sef14 antibody (lane 1) and anti-tag (T7) antibody (lane 2).

Figure 3 is a photograph showing results of a rSef14-latex agglutination assay for SE infection in

10

chickens exposed to S. enteritidis (A), S. pullorum (B), and serum-free antigen control (C).

Figure 4 is a photograph showing results of a rSef14-latex agglutination assay for SE infection in chickens exposed to S. enteritidis (A), S. gallinarum (B), S. pullorum (C), S. typhimurium (D), C. arizonae (E), E. coli (F), serum free antigen control (G), and serum control (H).

Figure 5 is a graph showing the percentage of chickens testing positive for anti-SE antibodies during 4 weeks post-innoculation. The five bars at each week represent innoculation with 10⁴, 10⁶, 10⁸, 10¹⁰, and control (no cells).

Figure 6 is a graph showing the antibody titres of chicken sera samples testing positive for anti-SE antibodies.

Figure 7 is a graph showing the antibody titres of chicken egg yolk samples testing positive for anti-SE antibodies.

20

25

30

15

Detailed Description of the Invention

The present invention is directed to a method for diagnosing Salmonella enteritidis infection or evidence of infection in an animal, particularly poultry, using a recombinant truncated fimbrial antigen.

"Infection" means active colonization of the animal by SE organisms. "Evidence of infection" means a prior history of colonization by SE in the animal, although active colonization is not present. Diagnosis of active infection is needed to protect against contamination of food supplies, whereas diagnosis of prior infection is

10

15

needed to alert against new infection or to trace the source of infection in a flock.

Fimbrial Proteins

Fimbriae are proteinaceous filamentous surface structures composed of protein subunits called fimbrin. These proteinaceous structures are thought to be virulence factors which mediate specific attachment to host cell mucosal surfaces. They are present in most enteric bacteria capable of invading host cells.

Salmonella enteritidis has four distinct fimbriae: Sef14, Sef17, Sef18 and Sef21 which are encoded by sefA, agfA, sefD and fimA genes, respectively. Sef14 is unique with only limited distribution in the genus. In contrast, all other fimbrial proteins are widely distributed in the genus. Thus, they have limited use as diagnostic reagents for SE detection.

Cloning and Expression of Sef14

In the present invention, a truncated form of the Sef14 antigen retaining the antigenic character of the entire protein has been produced. Unlike the complete protein, however, the truncated form can be easily produced in purified form and in large quantities, without special growth medium requirements.

PCR technology is used to produce the truncated Sef14 protein by amplification with suitable primers.

Primers are selected to amplify the gene encoding Sef14 in a region downstream of the encoded signal peptide, e.g., downstream of about nucleotide 145 of the DraI genomic fragment shown in Figure 1 of Turcotte and Woodward, Supra. Preferably, the PCR primers include

10

15

additional nucleotides at the 5' ends, encoding specific restriction enzyme recognition sequences, for ease of purification. For example, useful primers for amplifying that portion of the sefA gene encoding an immunogenic Sef14 fragment downstream of the signal peptide are shown below:

<u>GGGAATTC</u> GCTGGCTTTGTTGGTAACA		ID	NO:1
<u>GGGCTCGAG</u> TTAGTTTTGATACTGAACGTA	SEQ	ID	NO:2

After a truncated gene sequence encoding Sef14 is produced, it can be cloned into a host using a plasmid or phage as a vector. Typically, the expression of Sef14 fimbriae by cultured Salmonella enteritidis is highly dependent on the growth medium composition (Thorns et al, International Journal of Food Microbiology, 21:47-53 (1994)), and it is typically difficult to produce large quantities. However, a truncated form of Sef14 having at least the signal peptide removed is expressed in host systems such as E. coli without these difficulties.

Truncated Sef14 Antigen

Because the truncated Sef14 protein retains the antigenic characteristics of the complete protein, it is useful in various immunological methods. For example, the inventive antigen is useful in antibody binding immunoassays such as assays to detect the presence of antibodies against SE in a sample. Suitable binding assays include ELISA, wherein the recombinant Sef14 antigen is bound to a surface and exposed to antibodies against SE. To detect the presence of bound anti-SE antibodies, a marker such as an enzyme-linked secondary antibody is then added.

10

15

20

25

30

An agglutination assay using truncated Sef14 antigen-coated latex beads is preferred. In the agglutination reaction, antigen-coated latex beads form detectable clusters when exposed to antibodies against SE. This preferred assay is described more fully in Example 4, below.

Diagnostic Assays

The assays described above can be used to detect the presence of antibodies to Salmonella enteritidis.

Preferably, the assays are used to determine whether or not an animal, e.g. a poultry animal such as a chicken or turkey, is infected with SE. Animal fluid such as blood or serum can be used in a diagnostic assay. If an animal is infected with SE, the animal will typically produce anti-SE antibodies. The recombinant Sef14 antigen is used to detect the presence of anti-SE antibodies, SE infection or the SE organism itself. Diagnostic assays such as these are particularly useful in birds. More particularly, diagnostic assays are useful in detecting SE infections in chicken or turkey to prevent foodborne illness by poultry consumption.

Vaccine

Passive immunization with anti-Sef14 antibodies has been shown to reduce Salmonella enteritidis colonization (Peralta et al. 1994). Additionally, Sef14 can induce a T-cell immune response (Ogunniyi et al 1994). Because the truncated Sef14 antigen exhibits these immunological activities, can be produced in large quantities, and does not have the cumbersome growth requirements of the complete protein, the truncated Sef14

antigen is also useful as a vaccine to confer immunity against SE. Preferably, the truncated Sef14 antigen is used as a vaccine in poultry to prevent foodborne illnesses.

5

10

EXAMPLES

The invention may be better understood with reference to the following examples which are not intended to limit the invention.

Example 1

Isolation of S. enteritidis genomic DNA

S. enteritidis was grown overnight at 37°C in 15 Luria-Bertani (LB) broth. Genomic DNA was extracted as described (Sambrook, et al., 1989) using standard methods with minor modifications. In brief, bacterial cells were pelleted by centrifugation at 13,000 x g for 3 minutes, washed/suspended in 1 ml of 1 M NaCl, centrifuged for 5 20 minutes at 13,000 \times g, and the pellet resuspended in 1 ml TE buffer (50 mM Tris-HCl, 50 mM EDTA, pH 7.8). The sample was next incubated with 5 μ l of lysozyme (50 mg/ml) (Sigma Chemical Co., St. Louis, MO) and 0.3 mg/ml RNase A (Sigma) at 37°C for 30 minutes. To this suspension, 1% sarkosyl 25 and 0.6 mg/ml of proteinase K (Sigma) were added, and the mixture incubated at 37° for 1 hour. Following incubation, chromosomal DNA was extracted twice with an equal volume of phenol-chloroform-isoamyl alcohol (25:24:1) and once with chloroform:isoamyl alcohol (24:1). Genomic DNA in the 30 aqueous phase was precipitated at -20°C with two volumes of absolute ethanol and 0.1 volume of 3 M sodium acetate, and

10

15

pelleted by centrifugation at 13,000 x g for 5 minutes. The pellet was then washed twice with 70% ethanol, air dried, and suspended in TE Buffer (10mM Tris-HCl and 1mM EDTA pH 8.0). Total DNA was quantitated spectrophotometrically at A_{250} nm.

Example 2

Cloning of sefA gene fragment

Oligonucleotide primer selection and synthesis:

Oligonucleotide primers corresponding to an internal fragment (64-498 bp) of the open-reading frame of the sefA gene were used for PCR amplification. Additional bases were added to the 5' end of each primer in order to confer a recognition sequence for either EcoRI (forward primer) or XhoI (reverse primer). The oligonucleotide primers were obtained from Integrated DNA technologies Inc., Ames, IA. The DNA sequences for the forward and reverse primers are shown below:

<u>GGGAATTC</u> GCTGGCTTTGTTGGTAACA	SEQ	ID	NO:1
GGGCTCGAGTTAGTTTTGATACTGAACGTA	SEQ	ID	NO:2

20 Additional nucleotides added to the 5' end of the primers are underlined.

PCR amplification of sefA gene fragment.

Amplification reactions were performed in 30 μ l volumes with 30 pmol of each primer and 5 mM MgCl₂. The reagents and enzymes used for PCR were obtained either from Boehringer Mannheim (Indianapolis, IN) or Perkin Elmer (Foster City, CA). One hundred ng of genomic DNA was used as a template for PCR amplification with the following parameters: an initial denaturation at 94°C for 5 minutes, followed by 35 cycles of denaturation (94°C for 1.5

25

WO 98/03656 PCT/US97/12639

minutes), annealing 52°C for 1 minute) and extension (72°C for 2 minutes), and a final extension of 15 minutes at 72°C. All amplification reactions were performed in a Perkin-Elmer Cetus DNA thermal cycler (Model 480). The PCR products were analyzed on a 1% agarose gel, stained with ethidium bromide (0.5 μ g/ml), and photographed under UV light.

PCR products were gel extracted (Qiagen Inc., Chatsworth, CA), quantitated spectophometrically, at 260 nm, and cloned directly into pGEM-T vector (Promega, Madison, WI). Following ligation, 2μl of the reaction products were transformed into E. coli DH5α cells (Gibco BRL, Gaithersburg, MD) by the heat shock method. Recombinant colonies were selected on ampicillin/IPTG-Xgal containing plates and screened for the presence of the appropriate insert by restriction analysis.

Nucleotide sequence analysis

A bacterial colony containing the recombinant plasmid with the rSefA fragment was grown in LB-ampicillin media, and the plasmid extracted using Qiagen plasmid extraction kit (Qiagen). The nucleotide sequence of the insert was determined using oligonucleotide primers specific to the vector sequence by automated DNA sequencing at the University of Minnesota Advanced Genetic Analysis Center. The insert was sequenced in its entirety in both orientations, and the amino acid sequence deduced using the standard genetic code (DNA*, Madison, WI). Sequencing results are shown below for nucleotide and deduced amino acid sequences of the insert (Seq.ID.NO:5), together with a tag sequence added during the subcloning of the fragment into the pET/abc expression vector (Seq.ID.NO:3). The

5

10

15

20

25

added tag sequence at the 5' end, provides a Histidine-rich portion to facilitate purification of the sequence on nickel columns, as well as an antigenic region that specifically binds the T7 anti-tag antibody provided with the pET/abc vector kit.

Nucleic Acid Sequence encoding rSefA fragment SEQ ID NO:3

ATG	GGC	AGC	AGC	CAT	CAT	CAT	CAT	CAT	CAC	AGC	AGC	GGC	CTG	GTG	45	
CCG	CGC	GGC	AGC	CAT	ATG	GCT	AGC	ATG	ACT	GGT	GGA	CAG	CAA	ATG	90	
GGT	CGC	GGA	TGG	GAA	TTC	GCT	GGC	TTT	GTT	GGT	AAC	AAA	GCA	GTG	135	
GTT	CAG	GCA	GCG	GTT	ACT	ATT	GCA	GCT	CAG	AAT	ACA	ACA	TCA	GCC	180	
AAC	TGG	AGT	CAG	GAT	CCT	GGC	TTT	ACA	GGG	CCT	GCT	GTT	GCT	GCT	225	
GGT	CAG	AAA	GTT	GGT	ACT	CTC	AGC	ATT	ACT	GCT	ACT	GGT	CCA	CAT	270	
AAC	TCA	GTA	TCT	ATT	GCA	GGT	AAA	GGG	GCT	TCG	GTA	TCT	GGT	GGT	315	
GTA	GCC	ACT	GTC	CCG	TTC	GTT	GAT	GGA	CAA	GGA	CAG	CCT	GTT	TTC	360	
CGT	GGG	CGT	ATT	CAG	GGA	GCC	AAT	ATT	AAT	GAC	CAA	GCA	AAT	ACT	405	
GGA	ATT	GAC	GGG	CTT	GCA	GGT	TGG	CGA	GTT	GCC	AGC	TCT	CAA	GAA	450	
ACG	CTA	AAT	GTC	CCT	GTC	ACA	ACC	TTT	GGT	AAA	TCG	ACC	CTG	CCA	495	
GCA	GGT	ACT	TTC	ACT	GCG	ACC	TTC	TAC	GTT	CAG	CAG	TAT	CAA	AAC	540	
TAA	CTC	GAG	CCC	552	2											

*Additional amino acid residues added to the amino terminus to facilitate protein purification and cloning are underlined.

Deduced amino acid sequence of rSefA protein fragment*

Seq. ID NO: 4

MGSSHHHHHHSSGLVPRGSHMASMTGGOOMGRGSEFAGFVGNKAVVQAAVT
IAAQNTTSANWSQDPGFTGPAVAAGQKVGTLSITATGPHNSVSIAGKGASVSGG
VATVPFVDGQGQPVFRGRIQGANINDQANTGIDGLAGWRVASSQETLNVPVTT
FGKSTLPAGTFTATFYVQQYQN

*Additional amino acid residues added to the amino terminus to facilitate protein purification and cloning are underlined.

It is understood that the amino acids added to the N-terminus of the Spf14 antigen are optional, and used for ease of cloning and purification. The amino acid sequence in the absence of these added residues (Sequence ID No:6)

5

15

with or without other added residues for cloning or purification procedures, for example, are similarly useful as antigens in the diagnostic assays of the invention.

Subcloning sefA gene fragment into an expression vector

The pGEM-T plasmid carrying sefA fragment was double digested with EcoRI and XhoI, and the digested products gel purified (Qiagen) and cloned into EcoRI and XhoI digested pET/abc expression vectors (Novagen Inc., Madison, WI). Ligation products (2 µl) from each of the reactions were transformed into E. coli BL21(DE3)pLyS cells by heat shock method. Recombinant clones were cultured on kanamycin and chloramphenicol containing plates, and analyzed by restriction enzyme analysis.

15

20

25

30

10

5

rSefA fragment expression

The recombinant clones were selected based on restriction enzyme analysis with EcoRI and XhoI digestion, selecting those clones yielding appropriately sized fragments as compared with a vector control. Selected clones were analyzed for rSefA fragment expression. Briefly, a single colony from each (pETabc/SefA fragment) freshly streaked plate was picked and inoculated to 50 ml LB broth containing appropriate antibiotics and incubated with shaking at 200 rpm at 37°C until the OD₆₀₀ reached 0.6. Cultures were induced with IPTG (0.4 mM) and incubated for an additional 3 hours. Following incubation, the cells were pelleted and resuspended in 5 ml of TE buffer (50mM Tris-HCl pH 8.0, 2mM EDTA) and incubated with 25 µl of lysozyme (50 mg/ml) and 100 μ l of 1% Triton X-100 for 20 minutes at 30°C. The samples were sonicated until they were no longer viscous, and centrifuged at 39,000 x g for

10

15

20

25

30

20 minutes. The supernatant was passed through a 0.45 μm membrane filter, and stored at -20 $^{\circ}\text{C}$ until further use.

SDS-PAGE analysis

The cell lysates were next analyzed by SDS-PAGE for the presence of the rSefA fragment by mixing with an equal volume of 2x SDS solubilization buffer separating on 12% polyacrylamide gels, and staining with Coomassie blue. The results are shown in lane 1 of Figure 1 which contains the total protein produced by the vector and contained in the cell lysates.

Western blot analysis

The lysates were separated on 12% polyacrylamide gels and transferred onto a nitrocellulose membrane using Transblot apparatus (Bio-Rad laboratories, Hercules, CA). Following transfer, the membrane was blocked with 3% BSA in phosphate buffered saline (PBS) and stained with either T7 anti-tag antibody (Novagen) or rabbit anti-Sef14 specific antibody (kindly provided by Dr. W. W. Kay, University of Victoria, BC, Canada). The membrane was washed and stained with anti-rabbit IgG/HRP conjugate and treated with developing reagent (Amersham lif sciences, Inc., USA) for 1 minute, exposed to X-ray film, and the radiograph developed. The results are shown in Figure 2, where lane 1 is probed with anti-Sef14 antibody, and lane 2 with T7 anti-tag antibody.

Purification of rSef14 fragment protein by column chromatography and electroelution

The recombinant Sef14 protein fragment produced in the cell lysates described above was purified by binding

of the Histidine-rich tag to nickel columns as described by the manufacturer (Novagen). Briefly, the cells were induced and extract was prepared as described above except that the induced cells were suspended in Tris buffer The cell lysate was passed through nickel 5 without EDTA. columns and washed sequentially with binding buffer (5 mM imidazole, 0.5 M NaCl, 20 mM Tris-CHl, pH 7.9) and wash buffer (60 mM imidazole, 0.5 M NaCl, 20 mM Tris-HCl, pH The bound protein was eluted using elution buffer (1 10 M imidazole, 0.5 M NaCl, 20 mM Tris-HCl, pH 7.9), quantitated using a Bio-Rad protein assay kit (Bio-Rad laboratories, Hercules, CA), and analyzed by SDS-PAGE. (See Figure 1, lane 2, where the arrow indicates the rSef14 fragment at about 19 KDa.) Since the column purified 15 recombinant material contained traces of non-specific proteins, the appropriate rSef14 fragment was further purified by cutting the rSef14 fragment from the gel and electroelution (Bio-Rad) following the manufacturer's suggested protocol. The electroluted fragment is shown in 20 lane 4 of Figure 1 (at arrow).

Example 3

Covalent coupling of rSef14 to blue-dved latex beads

The electroeluted rSef14 protein fragment was coupled to either 0.5 μm or 1.0 μm blue-dyed latex beads (Polysciences Inc., Warrington, PA) by gluteraldehyde method. Briefly, 1 ml of 2.5% suspension of the beads were washed with PBS (pH 7.4), pelleted by centrifugation and resuspended in 1 ml of 8% gluteraldehyde (EM grade) in PBS, and incubated overnight with gentle end-to-end mixing at room temperature. Following gluteraldehyde treatment, the beads were pelleted, washed with PBS three times and

25

PCT/US97/12639

incubated with 500 µg of purified rSef14 fragment for 5 hours at room temperature with gentle end-to-end mixing. The beads were pelleted, and incubated with 1 ml of 0.5 M ethanolamine in PBS for 30 minutes at room temperature with gentle end-to-end mixing. The mixture was then treated with 1 ml of 10 mg/ml BSA in PBS for 30 minutes at room temperature, centrifuged and the pellet resuspended in 1 ml PBS (pH 7.4), containing 10 mg/ml BSA, 0.1% NaN₃ and 5% glycerol, and stored at 4°C to form rSef14 - fragment coated latex beads for use in agglutination assays.

Example 4

rSef14-latex bead agglutination test

Bacteria was administered to chickens by either injection, intratracheal or oral administration of 107 15 colony forming units (CFU) of either S. enteritidis, S. pullorum, S. arizonae, S. typhimurium, S. gallinarum, or E. coli. After about two to three weeks exposure, serum was collected and used to evaluate the sensitivity and specificity of the rSef14-latex beads in an agglutination 20 assay for anti-SE antibody binding. A total volume of 7.5 ul of rSef14 fragment coated latex beads, produced as described for Example 3, were mixed with an equal volume of chicken serum collected from birds exposed to various pathogens, as described above. The presence of 25 agglutination, visually seen as a loss of intense blue color in the sample (i.e., lightening of color as the coated beads agglutinate or form a lattice). Absence of the agglutination reaction was visualized by the remaining intense blue color of the dyed beads in a homogeneous 30 suspension. Positive or negative agglutination reaction

5

10

15

20

25

30

was recorded after two minutes. The results are shown in Figures 3 and 4.

In figure 3, intense blue color (negative result) is seen in test samples B and C (S. pullorum and the serum-free antigen control). In contrast, a positive agglutination result is seen in test Sample A, (S. enteritidis), as a pale blue, diffuse agglutination pattern.

In figure 4, a positive agglutination reaction is seen in sample A (S. enteritidis) and in sample H (serum control). No agglutination reaction is seen in the samples B-G containing serum animals exposed to the following pathogens: S. gallinarium (B), S. pullorum (C), S. typhimurium (D), S. arizonae (E), E. coli (F), and serum free antigen control (H).

Example 5

Detection of anti-S.E. anithodies in infected chickens

To confirm the specificity of the assay of the invention, forty SPF chickens (age 4weeks) were innoculated with various species of Salmonella. A suspension of 10° CFU in PBS was administered by injection. A booster dose of 10° CFU was administered orally two weeks later. Serum samples were taken at weekly intervals and assayed for the presence of anti-SE antibodies.

PCT/US97/12639

two assays are standard screening methods for the detection of Salmonella, using S. pullorium as a whole-cell antigen, and are not specific for SE, as shown in the table below.

To demonstrate the specificity of the assays of the invention, serum samples were assayed using the latex agglutination test (LAT) described above for Example 4, which utilized the truncated Spf14 antigen coupled to latex beads. Serum samples were also assayed for anti-SE antibodies by ELISA. In the ELISA, the truncated Spf14 antigen prepared as described for Example 3, was coated onto polystyrine plates. Antigen-coated plates were exposed to serum samples to permit binding of anti-SE antibodies to the antigen. The bound antigen-antibody complexes were washed, and then incubated with anti-chicken antibody coupled to biotin. The complex was then exposed to strep-avidin for signal detection.

Results are shown in the table below. The LAT and Elisa assays demonstrated a useful specificity for the detection of SE. Of the organisms tested, only S. dublin, a bovine pathogen, demonstrated cross-reactivity in the assays.

Species	SPT	MT	LAT	ELISA
S. enteritidis	+	+	. +	+
S. gallinarum	+	+	_	-
S. pullorum	+	+	-	_
S. dublin	+	+	+	+
S. berta	+	+	-	-
S. typhimurium	-	+	-	<u>-</u>
E. coli	-	-	-	<u> </u>
Control (no cells)	-	-	_	-

5

10

15

Example 6

Specificity of anti-SE assay

The ELISA assay for detecting anti-SE antibodies described above for Example 6 was tested for specificity using a panel of antisera against known pathogenic organisms. Each sera was assayed in the anti-SE ELISA. No crossreativity was observed with any of the tested antisera.

10

5

Antisera	ELISA	Antisera	ELISA
Pox	-	MG	-
Reo	_	NDV	. -
Rev	-	CAV	_
SB-1	-	HVT	-
IBDV	-	IBV	_
ILT	_	S.typhimurum	-
LLA	-	S.gallinarum	-
LLB	-	S.pullorum	-
MS			

15

Example 7

Sensitivity of ELISA for detection of SE

Fifty white leghorn layer chickens (5 weeks old) were orally innoculated in a single exposure with varied amounts of SE, from 10^4 to 10^{10} CFU in PBS. Serum samples were collected at weekly intervals for up to seven weeks. Eggs were collected for egg yolk antibody detection.

Samples were analized for detection of anti-SE

antibodies using the ELISA described above for Example 6. As shown in Figure 5, control chickens showed no positive reaction in the ELISA assay. Approximately 40-80% of chickens exposed to 10⁴, 10⁶, 10⁸, and 10¹⁰ CFU of SE tested positive for anti-SE antibodies during the first four weeks post-innoculation. From 4-7 weeks post-innoculation, the data stabilized at about 45% positive detection of anti-SE antibodies.

Antibody titers in the sera and egg yolks of chickens exposed to 10^4 , 10^6 , and 10^8 CFU of SE and testing positive in the ELISA for anti-SE antibodies are shown in Figures 6 and 7.

These data demonstrate specific detection of anti-SE antibodies using recombinant Sef14-antigen coated latex beads in an agglutination assay and using the antigen as a capture agent in an ELISA. These assays provide a sensitive and specific diagnostic tool for the detection of anti-SE antibodies in animals and for the diagnosis of SE infection.

5

10

20 SEQUENCE LISTING

- (1) GENERAL INFORMATION
- (i) APPLICANT: REGENTS OF THE UNIVERSITY OF MINNESOTA
- (ii) TITLE OF THE INVENTION:

 RECOMBINANT FIMBRIAL PROTEIN
- (iii) NUMBER OF SEQUENCES: 6
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Merchant, Gould, Smith, Edell, Welter & Schmidt
 - (B) STREET: 3100 Norwest Center, 90 South Seventh St
 - (C) CITY: Minneapolis
 - (D) STATE: MN
 - (E) COUNTRY: USA
 - (F) ZIP: 55402
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Diskette
 - (B) COMPUTER: IBM Compatible
 - (C) OPERATING SYSTEM: DOS
 - (D) SOFTWARE: FastSEQ Version 2.0
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE: 18-JUL-1997
 - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: 60/022,191
 - (B) FILING DATE: 19-JUL-1996
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Kettelberger, Denise M
 - (B) REGISTRATION NUMBER: 33,924
 - (C) REFERENCE/DOCKET NUMBER: 600.335W001
- (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 612/371-5268
 - (B) TELEFAX: 612/332-9081
 - (C) TELEX:

(2) INFORMATION FOR SEQ ID NO:1:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(ii) MOLECULE TYPE: cDNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
GGGAATTCGC TGGCTTTGTT GGTAACA	2
(2) INFORMATION FOR SEQ ID NO:2:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(ii) MOLECULE TYPE: cDNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
GGGCTCGAGT TAGTTTTGAT ACTGAACGTA	3
(2) INFORMATION FOR SEQ ID NO:3:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 552 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA 	
(ix) FEATURE:	
(A) NAME/KEY: Coding Sequence(B) LOCATION: 1540(D) OTHER INFORMATION:	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
ATG GGC AGC CAT CAT CAT CAT CAT CAC AGC AGC GGC CTG GTG CCG Met Gly Ser Ser His His His His His Ser Ser Gly Leu Val Pro 1 5 10 15	48
CGC GGC AGC CAT ATG GCT AGC ATG ACT GGT GGA CAG CAA ATG GGT CGC Arg Gly Ser His Met Ala Ser Met Thr Gly Gly Gln Gln Met Gly Arg 20 25 30	96

GGA Gly	TGG Trp	GAA Glu 35	TTC Phe	GCT Ala	GGC Gly	TTT Phe	GTT Val 40	GGT Gly	AAC Asn	AAA Lys	GCA Ala	GTG Val 45	GTT Val	CAG Gln	GCA Ala	144
GCG Ala	GTT Val 50	ACT Thr	ATT Ile	GCA Ala	GCT Ala	CAG Gln 55	AAT Asn	ACA Thr	ACA Thr	TCA Ser	GCC Ala 60	AAC Asn	TGG Trp	AGT Ser	CAG Gln	192
GAT Asp 65	CCT Pro	GGC Gly	TTT Phe	ACA Thr	GGG Gly 70	CCT Pro	GCT Ala	GTT Val	GCT Ala	GCT Ala 75	GGT Gly	CAG Gln	AAA Lys	GTT Val	GGT Gly 80	240
ACT Thr	CTC Leu	AGC Ser	ATT Ile	ACT Thr 85	GCT Ala	ACT Thr	GGT Gly	CCA Pro	CAT His 90	AAC Asn	TCA Ser	GTA Val	TCT Ser	ATT Ile 95	GCA Ala	288
	AAA Lys															336
	GGA Gly															384
ATT Ile	AAT Asn 130	GAC Asp	CAA Gln	GCA Ala	AAT Asn	ACT Thr 135	GGA Gly	ATT Ile	GAC Asp	GGG Gly	CTT Leu 140	GCA Ala	GGT Gly	TGG Trp	CGA Arg	432
	GCC Ala															480
AAA Lys	TCG Ser	ACC Thr	CTG Leu	CCA Pro 165	GCA Ala	GGT Gly	ACT Thr	TTC Phe	ACT Thr 170	GCG Ala	ACC Thr	TTC Phe	TAC Tyr	GTT Val 175	CAG Gln	528
	TAT Tyr			TAAC	TCGA	.GC C	C									552

(2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 180 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (v) FRAGMENT TYPE: internal
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Gly Ser Ser His His His His His Ser Ser Gly Leu Val Pro 10 Arg Gly Ser His Met Ala Ser Met Thr Gly Gly Gln Gln Met Gly Arg 25 Gly Trp Glu Phe Ala Gly Phe Val Gly Asn Lys Ala Val Val Gln Ala Ala Val Thr Ile Ala Ala Gln Asn Thr Thr Ser Ala Asn Trp Ser Gln Asp Pro Gly Phe Thr Gly Pro Ala Val Ala Gly Gln Lys Val Gly 70 Thr Leu Ser Ile Thr Ala Thr Gly Pro His Asn Ser Val Ser Ile Ala 90 Gly Lys Gly Ala Ser Val Ser Gly Gly Val Ala Thr Val Pro Phe Val 100 . 105 Asp Gly Gln Gly Gln Pro Val Phe Arg Gly Arg Ile Gln Gly Ala Asn 120 Ile Asn Asp Gln Ala Asn Thr Gly Ile Asp Gly Leu Ala Gly Trp Arq 135 140 Val Ala Ser Ser Gln Glu Thr Leu Asn Val Pro Val Thr Thr Phe Gly 150 155 Lys Ser Thr Leu Pro Ala Gly Thr Phe Thr Ala Thr Phe Tyr Val Gln 170 Gln Tyr Gln Asn

- (2) INFORMATION FOR SEQ ID NO:5:
- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 435 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Genomic DNA
- (ix) FEATURE:
 - (A) NAME/KEY: Coding Sequence
 - (B) LOCATION: 1...432
 - (D) OTHER INFORMATION:
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

GCT GGC TTT GTT GGT AAC AAA GCA GTG GTT CAG GCA GCG GTT ACT ATT

Ala Gly Phe Val Gly Asn Lys Ala Val Val Gln Ala Ala Val Thr Ile

1 5 10 15

GCA GCT CAG AAT ACA ACA TCA GCC AAC TGG AGT CAG GAT CCT GGC TTT 96
Ala Ala Gln Asn Thr Thr Ser Ala Asn Trp Ser Gln Asp Pro Gly Phe
20 25 30

ACA GGG CCT GCT GCT GCT GGT CAG AAA GTT GGT ACT CTC AGC ATT

Thr Gly Pro Ala Val Ala Ala Gly Gln Lys Val Gly Thr Leu Ser Ile

35

40

45

	GCT Ala 50									192
	GTA Val									240
	CCT Pro									288
	AAT Asn									336
	GAA Glu									384
	GCA Ala 130								T	433
7 7										

(2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 144 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (v) FRAGMENT TYPE: internal
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

10

25

*BNSDOCID: <WO ** 9803656A1*1A>

WE CLAIM:

1. A method for detecting anti- Salmonella enteritidis antibodies in animals, the method comprising:

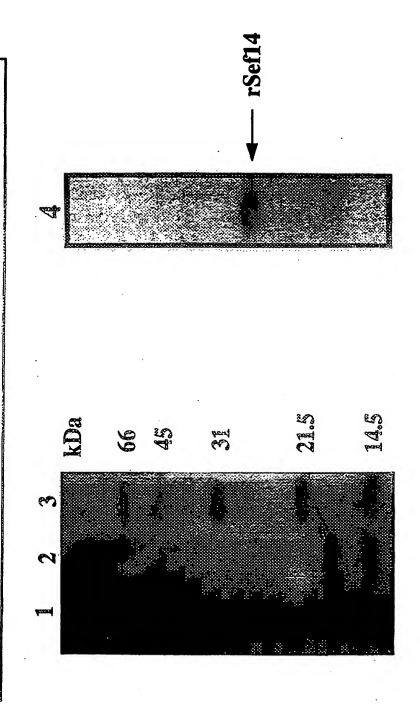
reacting a sample obtained from an animal with a truncated Sef14 antigen under conditions to permit anti-Sef14 antibodies to bind the antigen, the truncated antigen having at least the native Sef14 signal peptide removed; and

correlating antibody-antigen binding with the presence of anti-SE antibodies in the sample.

- 2. A method for diagnosing Salmonella enteritidis infection in animals, the method comprising:
- reacting a sample obtained from an animal with a truncated Sef14 antigen under conditions to permit anti-Sef14 antibodies to bind the antigen, the truncated antigen having at least the native Sef14 signal peptide removed; and
- 20 correlating antibody-antigen binding with Salmonella enteritidis infection.
 - 3. The method of claim 1, wherein the truncated Sef14 antigen has the amino acid sequence of Sequence ID. No. 4.
 - 4. The method of claim 1, wherein the truncated Sef14 antigen has the amino acid sequence of Sequence ID. No.6.
- 5. The method of claim 1, wherein said antigen is fixed to an inert surface prior to said reacting.

- 6. A Sef14 antigen consisting essentially of the amino acid sequence of Sequence I.D. No. 6.
- 7. An assay kit for the detection of anti-Salmonella enteritidis antibodies comprising an Sef14 antigen consisting essentially of the amino acid sequence of Sequence ID No. 6.
- 8. The assay kit of claim 7, wherein the antigen consists essentially of the amino acid sequence of Sequence ID. No. 4.
- 9. An antigen for stimulating the production of antiSalmonella enteritidis antibodies comprising the amino acid
 sequence of Sequence ID No. 4 or 6.
 - 10. The method of detecting anti-Salmonella enteritidis antibodies described in any of the foregoing claims, wherein the animal samples are obtained from fowl, and particularly from chickens or turkeys.

SDS-PAGE of rSell4 fragment protein



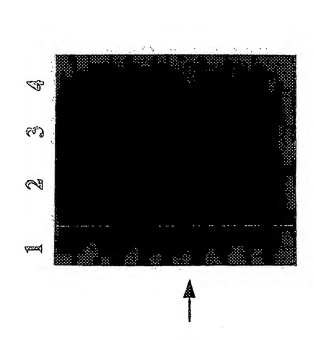
BL21DE3 (pET/sefA)before purification
 Purified rSef14 fragment protein

3. LMW marker 4. rSef14 after electroelution

FIG. 2

Western blot -- rSell4 probed with Sell4 monospecific polyclonal antibodies





Anti-Sef14
 T7 tag antibody
 BL21DE3 control
 LMW marker

Latex agglutination test

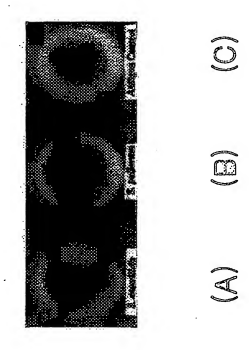
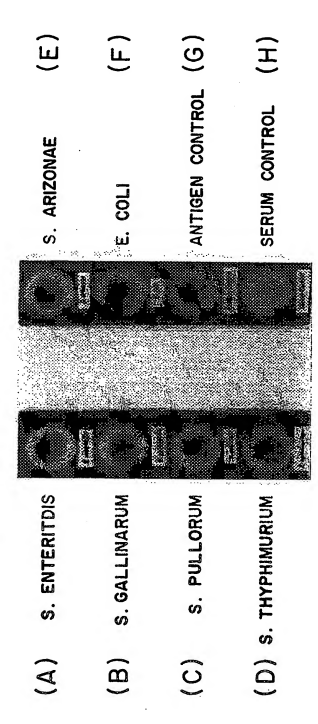
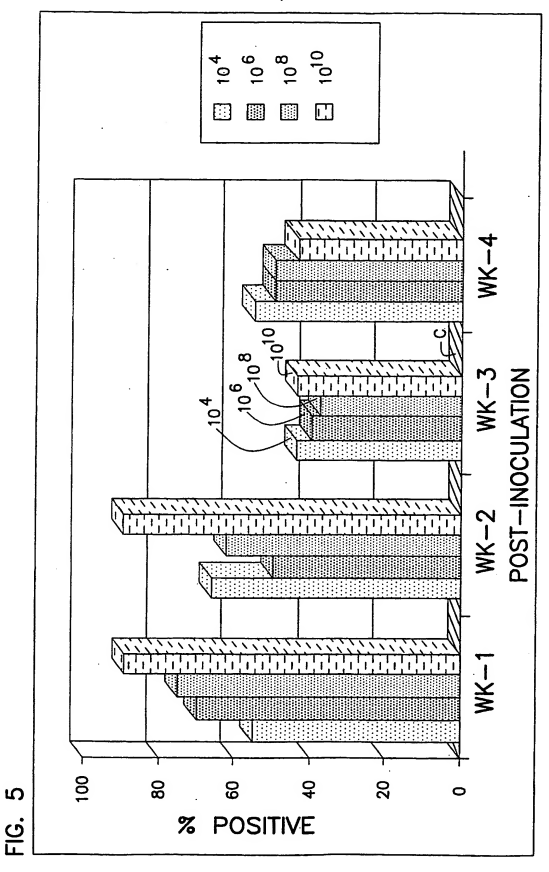


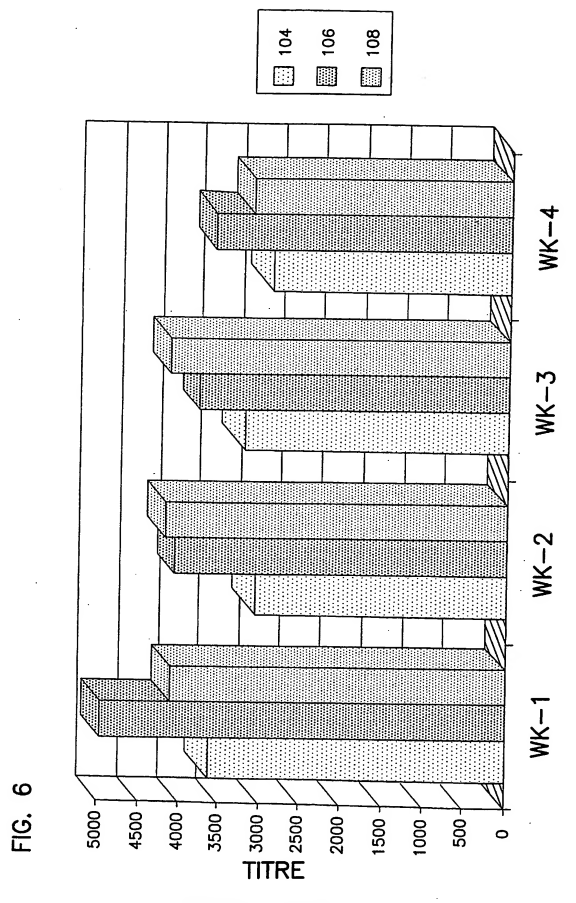
FIG. 4

Latex agglutination test



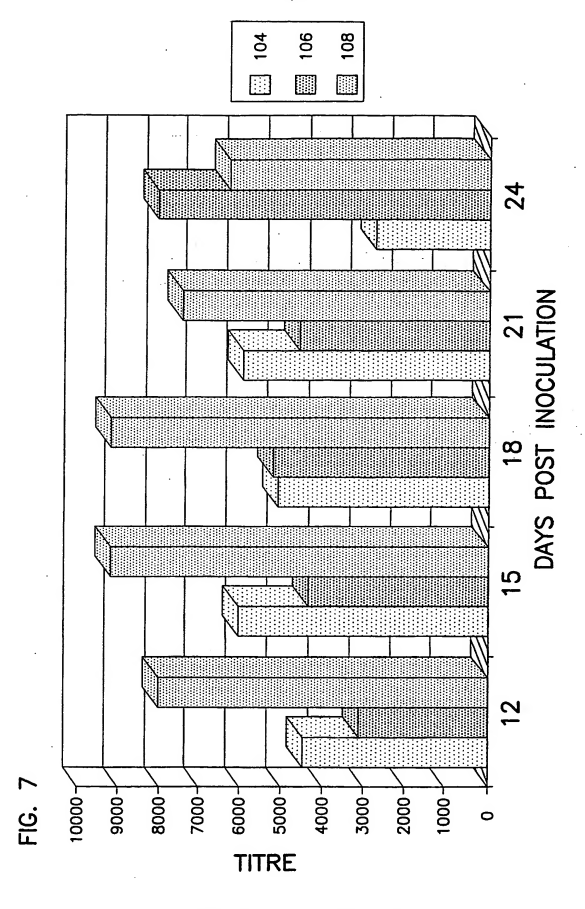


SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)





Inte .onal Application No PCT/US 97/12639

CLASSIFICATION OF SUBJECT MATTER A. CLASS IPC 6 C12N15/31 C07K14/255 G01N33/50 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 6 C12N C07K G01N Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category ° Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X THORNS CJ ET AL: "Development and 1,2,5,6, application of enzyme-linked immunosorbent assay for specific detection of Salmonella enteritidis infections in chickens based on antibodies to SEF14 fimbrial antigen." J CLIN MICROBIOL, APR 1996, 34 (4) P792-7, UNITED STATES, XP002047275 see page 792 - page 793; table 1 X WO 92 06197 A (MINI AGRICULTURE & 1,2,5,6, FISHERIES) 16 April 1992 see claims 2,3 Further documents are listed in the continuation of box C. X X Patent family members are listed in annex. Special categories of cited documents: *T* later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance invention earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-*O* document referring to an oral disclosure, use, exhibition or other means ments, such combination being obvious to a person skilled document published prior to the international filing date but later than the priority date claimed *&* document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 0 2. 12, 97 18 November 1997 Name and mailing address of the ISA Authorized offices European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Espen, J

Form PCT/ISA/210 (second sheet) (July 1992)

Interru al Application No
PCT/US 97/12639

	WALL A DOCUMENT OF THE PART OF THE PART	PC1/03 37/12033
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Category °	Cuation of document, with integration, where appropriate, or the relevant passages	Flansani (& Semi) 176.
Υ	THORNS CJ ET AL: "The use of latex particle agglutination to specifically detect Salmonella enteritidis." INT J FOOD MICROBIOL, JAN 1994, 21 (1-2) P47-53, NETHERLANDS, XP002047276 see the whole document	1,2,5,6, 10
Υ .	WO 92 06198 A (MINI AGRICULTURE & FISHERIES) 16 April 1992 see claims 1-28	1,2,5,6,
Υ	WO 93 20231 A (MINI AGRICULTURE & FISHERIES ;WOODWARD MARTIN JOHN (GB); THORNS CH) 14 October 1993 see claims 1-28	1,2,5,6, 10
A	CLOUTHIER SC ET AL: "Characterization of three fimbrial genes, sefABC, of Salmonella enteritidis." J BACTERIOL, MAY 1993, 175 (9) P2523-33, UNITED STATES, XP002047277	
	÷	

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

information on patent family members

Inter anal Application No
PCT/US 97/12639

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 9206197 A	16-04-92	AU 660152 B AU 8548991 A CA 2091982 A EP 0551325 A JP 6501934 T US 5510241 A AT 155169 T AU 660945 B AU 8656691 A CA 2091984 A DE 69126786 D EP 0551324 A WO 9206198 A	15-06-95 28-04-92 02-04-92 21-07-93 03-03-94 23-04-96 15-07-97 13-07-95 28-04-92 02-04-92 14-08-97 21-07-93 16-04-92
WO 9206198 A	16-04-92	AT 155169 T AU 660945 B AU 8656691 A CA 2091984 A DE 69126786 D EP 0551324 A JP 6502531 T AU 660152 B AU 8548991 A CA 2091982 A EP 0551325 A WO 9206197 A JP 6501934 T US 5510241 A	16-04-92 24-03-94
WO 9320231 A	14-10-93	AU 3895293 A	08-11-93